WO 2005/003161 PCT/EP2004/007046 FIGURE 1

HMMGENE FOR NT_006302.5

Peptide VMLMIQETNK (SEQ ID NO:18, frame = 1)
Predictions :
1223361-1223403 +

1224023-1224114 + VMLMIQETNK Score = 0.158

>NT_006302.5

MALSLSSSKRLQLDNRVMLMIQETNKQKVKGSGPYRNMTVTQMS*

FIGURE 2

HMMGENE FOR NT_006431.5

Peptide LLNNFPYR (SEQ ID NO:20, frame = 1)

Predictions: 2323834-2323931 + 2350894-2350927 + 2351807-2351866 +

2363588-2363659 + LLNNFPYR

Score = 0.019 >NT_006431.5

MISRQIGCRVYEDLRISKFFVISNCKDETAINGFYFGKNSIYGKVIEKTDEEIRSLFYE FPQTQLLNNFPYREVLWSLPVPRSSDRK* WO 2005/003161 PCT/EP2004/007046

FIGURE 3

HMMGENE FOR NT 007712.5

Peptide SGCTNLRSHQQCIR (SEQ ID NO:27, frame = 3)

Predictions: 116490-116500 + 121555-121669 + SGCTNLRSHQQCIR Score = 0.081 >NT_007712.5 MLASNSIFHFLRTLQTVLRSGCTNLRSHQQCIRVPFSPHPQ*

FIGURE 4

HMMGENE FOR NT 007914.5

Peptide ESIYFIIAAMLVATK (SEQ ID NO:31, frame = 1)

Predictions:

1365933-1366033 +

1374991-1375117 + ESIYFIIAAMLVATK

Score = 0.137

>NT_007914.5

MHRKDNGEMSAGEAGKAGTPKGEGHGKKPTHVISYSSSKRKSLFFWKESIYFIIAAMLV

ATKAANQIYEGQPTQS*

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FIGURE 5

HMMGENE FOR NT_009891.1

```
Peptide QADGTVFSK (SEQ ID NO:47, frame = 3)

Predictions:
995796-995822 +
1001710-1001787 + QADGTVFSK
Score = 0.080
>NT_009891.1

MIKTESKSKYLSFFTSFKQADGTVFSKMKRKHLK*
```

FIGURE 6

HMMGENE FOR NT_010909.5

```
Peptide CVIFPLNSYGMLLK (SEQ ID NO:53, frame = 3)
Predictions :
146623-146704 +
164675-164798 +
176327-176399 +
177711-177772 +
183840-183972 +
184830-184912 + CVIFPLNSYGMLLK
185665-185759 +
195917-196005 +
197742-198041 +
Score = 0.000
>NT 010909.5
MAFPDHKDAGKCSHLFSVPGEEREVKIGVPAVFCSCPCYVAELCCPILSRAPKPPDPVA
{\tt AEHLNHGQSRSDELSAYVSTYLVPGNVLGTGDPMTEDPTMERPYTFKDFLLRPRRDVSS}
{\tt ESDNNIRQINQEAAHRRFRSRRHISEDLEPEPSEGGDVPEIYYHENINLGEQK \underline{CVIFPL}
NSYGMLLKTISDQPSGAVRGTKQKASDHSRLQWGTVQLFDCWEERKDAKGRTYYVNHNN
RTTTWTRPIMQGAKDSPVRRAVKDTLSNPQSPQPSPYNSPKPQHKVTQSFLPPGWEMRI
{\tt APNGRPFFIDHNTKTTTWVRLLLFGSIFIMKSGINSLISLVFVVLAEELD} \\
```

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HMMGENE FOR NT 011896.6

FIGURE 8

HMMGENE FOR NT_019265.5

```
Peptide QVQSYHVLGK (SEQ ID NO:66, frame = 1)

Predictions:
745567-745650 +
749487-749540 + QVQSYHVLGK
Score = 0.014
>NT_019265.5

MAKPHIYPKYKNYLGVEALACGPTWKAEQVQSYHVLGKQRTNHIG*
```

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FIGURE 9

HMMGENE FOR NT_022148.5

```
Peptide CLCTHNGASKYMK (SEQ ID NO:73, frame = 1)

Predictions:
513123-513179 +
517522-517632 +
524582-524606 +
530347-530459 + CLCTHNGASKYMK
Score = 0.033
>NT_022148.5

MDLTLLETHLESYRISSQMPSFLLPLGQGGSTVIRDNVDPQKRAADLQESGQTIFQRKT
KTSEEGVNSPRRHNNPKCLCTHNGASKYMKQKHTEPDTSQLY*
```

FIGURE 10

HMMGENE FOR NT_022851.5

```
Peptide EQNKILSNLEIER (SEQ ID NO:76, frame = 3)

Predictions:
187415-187484 + EQNKILSNLEIER
191445-191529 +
202588-202654 +
203509-203661 +
204036-204245 +
Score = 0.023
>NT_022851.5

MEQDTKELKEQNKILSNLEIERDKEEAETQRNYEIFPRTCKCYELEPECKSRYQHLSEE
AEDMGLVVICPYLSEAAQSPQVFECIWSFLQISLVFISQNNLELVEISGKTLQDDYVTI
ARVICDQGGRVVNFGISWKLEVRGLDRDGKSCPQDPEKDSKEQPNLTEGEKAKGAVCKN
QISWSLASAKLLCVGRV*
```

FIGURE 11

HMMGENE FOR NT 007897.5

```
Peptide TEGLTLLQLV (SEQ ID NO:30, frame = 2)

Predictions:
1266436-1266537 + TEGLTLLQLV

Score = 0.108
>NT_007897.5

MDKRREAGNRESRISPGRVAGGRTEGLTLLQLV*
```

FIGURE 12

HMMGENE FOR NT_009561.5

```
Peptide MKPLVDYK (SEQ ID NO:43, frame = 1)

Predictions:
800421-800456 +
802726-802910 +
803769-803924 +
813981-814042 +
820146-820240 +
833001-833030 + MKPLVDYK
Score = 0.025
>NT_009561.5

MDMHRDKVYNEQGLIHMLFVAENEVSLFSSLVLCFLPMKRKKEKTPDGEPQIVGRFETP
LEFVLVMQSLVQTDNKTAFSDNFSYKSRLSDKLPSVPLPAWMHSWNLAFHKGIRIAFRQ
CFNHPKSRMYQSSLANTVLCASFDYLFRDEEPGLSNICTFSSSRLVQKVQLTATEGLLS
IRMKPLVDYK*
```

FIGURE 13

HMMGENE FOR NT_011387.5

```
Peptide DSYLNVKR (SEQ ID NO:55, frame = 2)

Predictions:
16190762-16190828 +
16193267-16193291 +
16197014-16197074 +
16205986-16206098 +
16210646-16210725 +
16212833-16212956 +
16214152-16214188 + DSYLNVKR
Score = 0.004
>NT_011387.5
MHNSPTVVTTQYSLTDEWIIKWVMIYQRNQGNNCSRGSGFTFWLGDYKHSVDPSIASPS
PEAAALCVPDDNLGIGTNQYQEWVCWERALRLTRMDSINQAPLPCILSCIGAMEATALL
RPVSCLTFRKCVDYFWLRVEREIAWERKSSYECQLNFGCFYKDSYLNVKR*
```

FIGURE 14

HMMGENE FOR NT_027064.2

```
Peptide PEMVVQACSLSY (SEQ ID NO:96, frame = 3)

Predictions:
553561-553602 +
578259-578297 + PEMVVQACSLSY
Score = 0.027
>NT_027064.2
MGKGWEVYNRQDLQPEMVVQACSLSY*
```

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FIGURE 15

HMMGENE FOR NT_028428.2

Peptide NAKISQAPW (SEQ ID NO:102, frame = 1)

Predictions : 290705-290780 + 296029-296147 + NAKISQAPW Score = 0.091>NT_028428.2

MDASVGHYPKKINTGMENQVPHVLASLWEAEVGESPEVRSSKPDWPRWQNPISTKNAKI

SQAPW*

FIGURE 16

HMMGENE FOR NT_019546.5

Peptide NPMKIFEK (SEQ ID NO:67, frame = 1)

Predictions: 611472-611544 + NPMKIFEK 612968-613179 + Score = 0.475>NT 019546.5

MMLRNPMKIFEKRKYIPQQKMLQK*